(1) Publication number:

0 376 448 Δ1

(12)

EUROPEAN PATENT APPLICATION

(21) Application number: 89311296.1

(51) Int. Cl.5: A61L 9/01

② Date of filing: 01.11.89

(3) Priority: 29.12.88 JP 334932/88 29.06.89 JP 169351/88

- (43) Date of publication of application: 04.07.90 Bulletin 90/27
- Designated Contracting States: AT BE CH DE ES FR GB IT LI NL SE
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- (54) Deodorizer composition.
- (57) A deodorizer composition comprising:
 - (a) an activated charcoal, a silica gel, a clay mineral, or an aluminosilicate having the following composition, in terms of oxides, represented by the composition ratio of the three components:

SiO₂: 5 to 80 mole%;

 $Mo_{n/2}$: 5 to 65 mole%; Al₂O₃: 1 to 60 mole%

wherein M represents at least one metal selected from the group consisting of zinc, copper, silver, cobalt, nickel, iron, titanium, barium, tin and zirconium, and n represents a valence of metal; and

(b) at least one component selected from the group consisting of oxidizing agents, plant extracts, and germicides.

DEODORIZER COMPOSITION

BACKGROUND OF THE INVENTION

1. Field of the Invention

The present invention relates to a deodorizer composition, more specifically, to a deodorizer composition having a strong deodorizing power which can be widely utilized to eliminate objectionable odors in the home and mitigate objectionable industrial odors generated by factories and the like.

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2. Description of the Related Art

The increasing urbantization of formerly sparsely-populated areas has brought more people into daily contact with the various smells and odors of their environment, and this has led to a strong interest in, and severe criticism of, particularly offensive odors. Various methods of treating offensive odors are known in the art, of which the following are typical:

- (1) Sensitive Deodorizing ... masking with perfumes or flavors;
- (2) Physical Deodorizing ... adsorption with, for example, activated carbon or absorption or clathrating with, for example, cyclodextrin;
- (3) Chemical Deodorizing ... neutralization with an acid or alkali, oxidation or reduction with an oxidizing agent or reducing agent, or an addition of, for example, lauryl methacrate; and
- (4) Biological Deodorizing ... deodorizing by germicidal action with germicide, or effects obtained with microorganisms or enzymes.

The sensitive deodorizing method is often unsuccessful since only the flavor of the odor is changed and the offensive odor itself is still present although masked, and thus if the balance between the offensive odor and the flavor is lost, the offensive odor again prevails and the balance must be restored.

In the physical deodorizing method, although the offensive odor is absorped or clathrated, problems arise in that the adsorption power or clathrating power is not strong enough for practical use.

In the chemical deodorizing method, certain chemicals should not be used from the viewpoint of safety, and although a chemical deodorization of one offensive odor can be achieved, this same process will have no affect on the other various offensive odors generated in daily life.

In the biological deodorizing method, disadvantages in the deodorizing rate and a continuous effect or durability arise and, therefore, universal effects cannot be obtained by a single deodorizing method.

Conventionally activated carbon is most widely used, and is known as a deodorizer capable of adsorbing various offensive odor components. Nevertheless, among such offensive odor components, the capability of activated carbon for the adsorption of lower amines is small, and in particular, the deodorizing power thereof against hydrogen sulfide and ammonia is low.

Various attempts have been made to solve the above-mentioned problems. For example, JP-A-55-51421 (i.e., Japanese Unexamined Patent Publication) proposes that halides be supported on activated carbon:

JP-A-53-137089 proposes that metals be supported on activated carbon; and the adhesion or deposition of acids or alkalis has been studied. Nevertheless, these proposals are not still satisfactory.

Furthermore, although the above-mentioned patent publications generally describe deodorizing performances against various objectionable or offensive odors, the deodorizing speed (or rate) and the deodorizing amounts are also important factors, taking into consideration the utility of the deodorizer compositions.

The present inventors have proposed, in JP-A-63-220874, the use of aluminosilicates having a specified SiO_2 , $MO_{n/2}$ and Al_2O_3 composition in terms of the oxides expressed by the three component composition as a deodorizer.

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SUMMARY OF THE INVENTION

Accordingly, the objects of the present invention are to eliminate the above-mentioned disadvantages of the prior art and to provide a deodorizer composition exhibiting excellent deodorizing effects for various objectionable odors, and having a high deodorizing speed and safety.

Another object of the present invention is to provide a deodorizer composition exhibiting an excellent deodorizing effect for various objectionable odors, and an excellent persistency of the effect.

Other objects and advantages of the present invention will be apparent from the following description.

In accordance with the present invention, there is provided a deodorizer composition comprising:

(a) an activated charcoal, a silica gel, a clay mineral, or an aluminosilicate having the following composition, in terms of oxides, represented by the three component composition ratios:

SiO₂: 5 to 80 mole%;

Monn: 5 to 65 mole%;

Al₂O₃: 1 to 60 mole%

wherein M represents at least one metal selected from the group consisting of zinc, copper, silver, cobalt, nickel, iron, titanium, barium, tin and zirconium, and n represents a valence of metal; and

(b) at least one component selected from the group consisting of oxidizing agents, plant extracts, and germicides.

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DESCRIPTION OF THE PREFERRED EMBODIMENT

The activated charcoal, silica gel, clay mineral, and aluminosilicate usable as the component (a) in the present invention preferably has a BET specific area of $100 \text{ m}^2/\text{g}$ or more, more preferably 500 to 2000 m^2/g , when determined with N_2 gas.

Activated charcoal may be obtained by a treatment of, for example, coal, petroleum residue, charcoal, fruit, and shells, by the gas activation method using, for example, steam, carbon dioxide, or the chemical activation method using, for example, zinc chloride.

As the aluminosilicate, zeolite and those having the composition ratio of the three components represented in terms of oxides shown below are preferable:

SiO₂: 5 to 80 mole%, preferably 25 to 75 mole%;

Mo_{n/2}: 5 to 65 mole%, preferably 15 to 60 mole%;

Al₂O₃: 1 to 60 mole%, preferably 1 to 45 mole%

wherein M represents at least one metal selected from zinc, copper, silver, cobalt, nickel, iron, titanium, barium, tin and zirconium, and n represents a valence of metal.

Of the component (a), aluminosilicates, in particular are considered to have the characteristics of a solid acid and solid acid group, and it has been reported that the acidity varies depending on the content of alumina in silica-alumina type catalysts, and that the acidity differs depending on the ratio of the contents of alumina and silica (Kozo Okabe: Acidic and Basic Catalysts, p. 183 (1967)). Further, since the aluminosilicate has a structure in which acidic SiO₂ and basic metal oxides are bound together, it has basic and acidic polarities and may be considered to exhibit deodorizing effect for objectionable odor components based primarily on a chemical adsorption and a chemical reaction.

The aluminosilicate is obtained as a white or pale-colored powder, and is prepared by allowing a water soluble silicate, a water-soluble metal salt, and further, a water-soluble salt aluminum salt and/or a water-soluble aluminate, in amounts corresponding to the above-mentioned composition ratio, to react in the presence of water, and if necessary, heating the precipitates obtained in the presence of water.

The above-mentioned reaction proceeds steadily due to a metathetical reaction. Namely, when an alkali silicate such as sodium silicate is used as a silica component, a water-soluble metal salt such as a chloride, nitrate, and sulfate is used as a metal oxide component, and further, sodium aluminate and/or a water-soluble aluminum salt such as aluminum chloride or aluminum sulfate are used as the alumina component, these components are mixed together in the presence of water, followed by effecting the metathetical reaction.

To uniformly effect the above-mentioned metathetical reactions, it is preferable to simultaneously add, to an aqueous dispersion containing silica previously dispersed therein, an aqueous silicate solution, an aqueous metal salt solution and an aqueous solution containing the alumina component, while allowing a reaction.

The above-mentioned metathetical reaction can be carried out at room temperature but, of course, also can be carried out upon heating at a temperature of up to, for example, about 95 °C.

The preferable pH range of the reaction during the simultaneous addition is 5 to 10, most preferably especially 6 to 9. If necessary, the pH of the reaction mixture can be controlled by the addition of an acid or alkali, to maintain the above-mentioned pH range.

The above-mentioned simultaneous addition of the reactants allows the formation of precipitated aluminosilicate having a composition substantially corresponding to that of the aqueous solution. The

precipitates thus formed are separated, or optionally, are heated in the presence of water, to obtain a white or pale-colored fine powder.

As the oxidizing agent (i) of the component (b), chlorine dioxide, sodium hypochlorite, calcium hypochlorite, sodium dichloroisocyanurate, sodium p-chlorotoluenesulfonchloroamide, hydrogen, peroxide, sodium percarbonate, potassium persulfate, potassium monopersulfate, sodium persulfate, ammonium persulfate, potassium permanganate, diperoxydodecane diacid, and magnesium perfumarate, are included as representative examples. Among the above, percarbonates, persulfates such as sodium persulfate, potassium persulfate, ammonium persulfate, and potassium monopersulfate; organic peracids such as diperoxydodecane diacid, and magnesium perfumarate, and salts thereof, are preferred.

As the plant extract (ii) of the component (b), an extract of a plant selected from red algae, brown algae, gymnosperms and angiosperms may be employed, preferably one having a deodorizing effect. Examples of such plants are as follows:

1) Rhodophyta

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- i) Gigartinales
- a) Gracilariaceae Gracilaria verrucosa Papenfuss
 - b) Gigartinaceae Chondrus ocellatus Halmes
 - ii) Ceramiales
 - a) Rhodomelaceae L. Pinnata, L. Intermedia
 - 2) Phaeophyta
 - i) Chordariales
 - a) Chordariaceae Nemacystus decipiens Kuckuck
 - ii) Laminariales
 - a) Laminariaceae Eisenia bicyclis Setchell
 - iii) Fucales
- a) Sargassaceae Hizikia fusiforme Okamura, Sargassum fulvellum C. Agardh, Sargassum horneri C. Agardh, Sargassum serratifolium C. Agardh, Sargassum thunbergii 0. Kuntze, Sargassum serratifolium C. Agardh, Sargassum ringgoldianum Harvey, Sargassum patens C. Agardh, Sargassum tortile C. Agardh, Sargassum hemiphyllum C. Agardh,
 - iv) Laminariales
- 30 a) Laminariaceae Undaria pinnatifida Suringar
 - 3) Gymnospermae
 - i) Ginkgoales
 - a) Ginkgoaceae Ginkyo biloba L.
 - ii) Pinales
- s a) Taxodiaceae Metasequia
 - 4) Angiospermae
 - i) Piperales
 - a) Piperaceae Piper nigrum L, Cubebs
 - ii) Fagales
- a) Betulaceae Betula playtphylla Sakatchev var japonica Ha
 - b) Fagaceae Quercus
 - iii) Polygonales
 - a) Polygonaceae Rheum undulatum L., Polygonum Hydropiper L
 - iv) Ranales
- a) Magnoliaceae Magnolia Kobus DC, Magnolia oborata Thunb
 - b) Ranunculaceae Paeonia suffruticosa Andr., Coptis japonica Makino, Cimicifuga simplex Wormsk. var. ramosa Maxim.
 - v) Ranales
 - a) Lardizabalaceae Akebia quinata Decne
- b) Berberidaceae Nandina domestica Thunb, Epimedium macranthum Morr. et Decne. Var. violaceu
 Franch
 - c) Lauraceae Cinnamonum camphora Sieb
 - vi) Rhoeadales
 - a) Papaveraceae Papaver somniferum L.
- 55 b) Cruciferae Isatis indigotica Fortune
 - vii) Sarraceniales
 - a) Droseraceae Drosera rotundifolia L
 - viii) Rosales

- a) Hamamelidaceae Witch hazel
- b) Rosaceae Prunus donarium Steb, Duchesnea chrysantha Miquel, Crataegus cuneata Sneb. et Zucc
- c) Leguminosae Cercis chinensis Bunge, Hedysarum esculentum Ledef
 - ix) Geraniales
- a) Rutaceae Xanthoxylum piperitum DC., Phellodendron amurense Rupr.
 - b) Meliaceae Melia azedarach L. var. japonica Makino
 - x) Parietales
 - a) Theaceae Thea sinensis L., Camellia japonica L.
 - xi) Myrtiflorae
- a) Punicaceae Punica grantum L.
 - b) Myrtaceae Syzygium aromaticum, Eucalyptus globulus Labill, Myrtaceae pimenta officinalis
 - xii) Umbelliflorae
 - a) Umbelliferae Apium graveolens L. Anise pimpinella anisum, Nothosmyrnium japonicum Miq xiii) Ericales
- a) Ericaceae Pieris japanica D. Don.
 - xiv) Contortae
 - a) Oleaceae Forsythia syspensa Vahl.
 - xv) Tubiflorae
 - a) Labiatae Salvia officinalis L., Thymus, vulgasis, Origanum spp., Rosmarinus officinalis L., Skull-cap, Elsholtzia ciliata Hylander, Leonurus sibiricus L., Mentha spicata L. var. crispa Benth, Mosla dianthre Maxim, Perilla frutescens Britton Var. acuta Kudo, Ocymum basilicum L., Origanum vulgare, Satureia ssp.
 - b) Solanaceae Lycium chinense Mill
 - c) Scrophulariaceae Picrorrhiza Kurroa Rogl.,
 - xvi) Rubiales
 - a) Rubiaceae Uncaria gambir Roxb.
 - xvii) Campanulatae
 - a) Compositae Chicory, Arnica

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- xviii) Scitamineae
- a) Zingiberaceae Curcuma zedoaria Roscoe, Hedy chium spicatum Hamilton

To obtain solvent extracts from these plants, as the plant, either the whole grass or the respective sections of plants such as leaves, plant skin, flowers, fruit skin, fruit, root stalk, or root may be employed, and the section containing relatively more of the deodorizing effective ingredient of the plant can be selected.

When obtaining deodorizing effective ingredients from these plants, known methods can be employed. For example, it is possible to employ the method in which plants dried and then cut and formed into powder, are extracted with at least one polar solvent such as water, ethyl ether, ethylene chloride, dioxane, acetone, ethanol, n-butanol, ethyl acetate, propylene glycol or at least one nonpolar solvent such as nhexane, petroleum ether, ligroin, cyclohexane, carbon tetra chloride, chloroform, dichloromethane, 1,2dichloroethane, toluene, benzene, or a mixed solvent of these polar solvents and nonpolar solvents. In this case, as the extraction operation, a conventional method can be employed in which, for example, plants are subjected to maceration in a solvent.

The extract obtained by the extraction operation as described above is usually formulated in a deodorizer after evaporation of the solvent, but in some cases, particularly when the solvent is water, ethanol can be also formulated as such in a deodorizer without evaporation of the solvent.

Examples of the germicide (iii) of the component (c) include p-oxybenzoate, o-phenylpheol, 3-methyl-4isopropylpheol, bistributyltin oxide, dibutyltin naphthenate, triphenyltin oxide, benzetonium chloride, benzalkonium chloride, α-bromocinnamic aldehyde, alkyl-di(aminoethyl)glycine salt, 2-bromo-2-nitropropane-1,3diol, chlorohexidine, 2-(4-thiazolyl)benzoimidazole, stearyl dimethylbenzylammonium chloride, distearyl dimethylammonium chloride, stearyl trimethylammonium chloride, cetylpyridium chloride, domiphen chloride, alkylisoquinolium bromide, 3,4,4 trichlorocarbanilide, dehydroacetic acid, sorbic acid, p-chloro-mxylenol, p-chloro-m-cresol, 2,4,6-tribromophenol, 2,4,6-trichlorophenol, pentachlorophenol, 2,4,4'-trichloro-2'hydroxydiphenyl ether, 3-trifluoromethyl-4,4'-dichloro-N,N'-diphenylurea, dimethyldithiocarbamate, 1,2benzisothiazolin-3-one, butyl-m-cresol, a-chlorobenzaldoxime acetate, resorcin, isopropylmethylphenol, and resorcin monoacetate. Among the above, p-chloro-m-xylenol, butyl-n-cresol, o-phenylpheol, α -bromocinnamic aldehyde are preferred.

The deodorizer composition of the present invention can be obtained with the above components (a) and (b) as the essential components.

These respective components may be formed into powdery or granular products and mixed, the

component (a) may be impregnated with the component (b), and a carrier may be impregnated or attached with the component (b) and mixed with the component (a), or further, the components (a) and (b) may be kneaded and granulated under the condition maintained in a solvent.

In the deodorizer composition of the present invention, it is preferable to contain the component (a) and the component (b) at a weight ratio of (a)/(b) = 1/99 to 99.99/0.01, more preferably 50/50 to 99.5/0.5. Also, if necessary, optional components can be further formulated in the composition.

In the composition of the present invention, in addition to the components (a) and (b), optional components such as surfactants, alkali agents, acid substances, dyes, UV-ray absorbers, antioxidants, and polymeric substances can be also formulated, if necessary.

A gas having an objectionable odor generally consists of a large number of components. As representative objectionable odor components, nitrogen type gases such as ammonia and amine, and sulfur type gases such as hydrogen sulfide and mercaptans, are known and are generated in large amounts. Nevertheless, since these nitrogen type and sulfur type objectionable odor components differ in behavior, an effective deodorizing base material for both objectionable odor components was not known.

In the present invention, by using the component (a) and the component (b), it is now possible to deodorize various objectionable odors. The performances demanded of deodorizers include, in addition to effects against various objectionable odors, a rapid deodorizing speed and large deodorizing capacity, i.e., long-term persistency.

In the present invention, the component (a), provides a deodorizing effect against various objectionable odors, and further, by using the component (b) in combination therewith, the deodorizing speed and the deodorizing capacity are improved.

Of the porous substances of the component (a), aluminosilicates in particular are considered to have the characteristics of a solid acid and solid acid group, and it has been reported that the acidity varies depending on the content of alumina in silica-alumina type catalysts, and that the acidity differs depending on the ratio of the contents of alumina and silica [Kozo Okabe: Acid Base Catalyst, p. 183 (1967)]. Further, since the aluminosilicate has a structure in which acidic SiO₂ and basic metal oxides are bound together, it has basic and acidic polarities and is considered to exhibit a deodorizing effect against objectionable odor components based primarily on a chemical adsorption and chemical reaction.

On the other hand, porous substances having a specific surface area of 100 m²/g or more as represented by activated charcoal have an excellent physical adsorption.

The deodorizing mechanism in the deodorizer composition of the present invention has not been defined, but it is considered that the physical adsorption proceeds at a very rapid speed until reaching an equilibrium, and subsequently, a chemical reaction occurs. That is, by combining a porous substance of the component (a) and the component (b), a rapid adsorption of objectionable odor onto a porous substance with a large specific area, and a chemical reaction of the adsorbed objectionable odor with the component (b), are considered to act synergetically, thereby effectively supplying a stronger deodorization effect.

Also, the objectionable odor trapped by the adsorption and chemical reaction will not be released as an odor, and it is considered that a release of an odor, which is a drawback of a deodorization by physical adsorption, is inhibited to improve the durability of the deodorizing effect.

According to the deodorizer composition of the present invention, by using the component (a) and an oxidizing agent (i) as the component (b) in combination, a stronger deodorizing power can be exhibited than in the case where each is used alone, but the deodorizing effect can be exhibited more rapidly and the persistency of the effect is prolonged.

In the present invention, by using the component (a), a plant extract (ii), and/or a germicide (iii) as the component (b) in combination, it is now possible to deodorize various objectionable odors.

Also, by using a plant extract (ii) of the component (b) in combination, in addition to eliminating various objectionable odors, a deodorizer composition is realized which has a rapid deodorizing speed, large deodorizing capacity, and a high safety factor.

Further, by combining a germicide (iii) of the component (b) with the component (a), the deodorizing speed and the deodorizing capacity are improved to provide a deodorizer having a prolonged persistency.

Therefore, although various objectionable odor components are exuded from the human body, in the home, and by industry facilities, the deodorizer composition of the present invention rapidly eliminates these objectionable odors and provides a long-term deodorizing effect, and thus can be widely utilized.

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EXAMPLE

The present invention will now be further, illustrated by, but is by no means limited to, the following synthetic Examples of the alumino silicates, Examples of the deodorizer compositions, and results of the evaluation thereof.

Synthetic Example 1

A 109 g amount of #3 sodium silicate (SiO₂: 22.0%, Na₂O: 7.0%) and 94 g of sodium hydroxide (NaOH content: 2.35 mole) were dissolved in water to prepare 1 liter of a solution A (SiO₂ content: 0.4 mole). Further, 95 g of zinc chloride (anhydrous salt) and 97 g of aluminum chloride (hexahydrate) were dissolved in water to prepare 1 liter of a solution B (ZnO content: 0.7 mole, Al₂O₃: 0.2 mole).

Then, one liter of water was charged into a 5 liter beaker and, while stirring, the solutions A and B were simultaneously added thereto at a feed rate of about 25 cc/min, respectively. The pH of the resultant reaction mixture was 6.9 after the addition.

The stirring was further continued, and after aging for 30 minutes, the mixture was heated at a temperature of 85 to 90°C for 2 hours on a water bath. The reaction mixture thus obtained was filtered with aspiration, followed by washing with water and drying at a temperature of 110°C, and the cake thus obtained was sifted through a sieve to obtain zinc aluminosilicate in the form of white particles having a size of 8 to 16 mesh (Tylor).

The composition of the three components and the BET specific surface of the resultant particles are shown in Table 2, together with the data of the following Synthetic Examples.

Synthetic Example 2

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In the same manner as in Synthetic Example 1, 139 g of #3 sodium silicate (SiO₂: 22.0%, Na₂O: 7.0%) and 88 g of sodium hydroxide (NaOH content: 2.2 mole) were dissolved in water to prepare 1 liter of a solution A (SiO₂ content: 0.51 mole). Further, 65 g of zinc chloride (anhydrous salt) and 126 g of aluminum chloride (hexahydrate) were dissolved in water to prepare 1 liter of a solution B (ZnO content: 0.48 mole, Al₂O₃: 0.26 mole).

Then, one liter of water was charged into a 5 liter beaker and, while stirring, the solutions A and B were simultaneously added thereto at a feed rate of about 25 cc/min, respectively. The pH of the resultant reaction mixture was 8.6 after the addition.

The reaction mixture obtained above was treated in the same manner as in Synthetic Example 1, whereby zinc aluminum silicate in the form of particles having a size of 8 to 16 mesh was obtained.

Synthetic Example 3

A 273 g amount of #3 sodium silicate (SiO₂: 22.0%, Na₂O: 7.0%) and 60 g of sodium hydroxide (NaOH content: 1.5 mole) were dissolved in water to prepare 1 liter of a solution A (SiO₂ content: 1.0 mole). Further, 34 g of silver nitrate and 225 g of aluminum nitrate (nonahydrate) were dissolved in water to prepare 1 liter of a solution B (Ag₂O content: 0.1 mole, Al content: 0.3 mole).

Then, one liter of water was charged into a 5 liter beaker and, while stirring, the solutions A and B were simultaneously added thereto at a feed rate of about 25 cc/min, respectively. The pH of the resultant reaction mixture was 8.9 after the addition.

The stirring was further continued, and after aging for 1 hour, the cake thus obtained was sifted through a sieve to obtain silver aluminosilicate in the form of particles having a size of 8 to 16 mesh.

Synthetic Example 4

A 77 g amount of #1 sodium silicate (SiO₂: 35.0%, Na₂O: 17.5%) and 24 g of sodium hydroxide (NaOH content: 0.6 mole) were dissolved in water to prepare 1 liter of a solution A (SiO₂ content: 0.45 mole). Further, 216 g of zinc sulfate (heptahydrate) and 75 g of sodium aluminate (Al₂O₃ content: 20.5%, Na₂O content: 19.5%) were dissolved in water to prepare 1 liter of a solution B (ZnO content: 0.75 mole, Al₂O₃: 0.15 mole).

Then, one liter of water was charged into a 5 liter beaker and, while stirring, the liquids A and B were

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simultaneously added thereto at a feed rate of about 25 cc/min, respectively. The pH of the resultant reaction mixture was 6.9 after the addition.

The stirring was further continued, and after aging for 30 minutes, the mixture was heated at a temperature of 85 to 90 °C for 2 hours on a water bath. The reaction mixture thus obtained was filtered with aspiration, followed by washing with water and drying at a temperature of 110 °C, and the cake thus obtained was sifted through a sieve to obtain zinc aluminosilicate in the form of white particles having a size of 8 to 16 mesh.

Synthetic Examples 5 to 12

In the same manner as in Synthetic Example 3 (silver salt), aluminosilicates containing cobalt, nickel, iron, copper, titanium, barium, tin, and zirconium were obtained from the solutions A and B listed in Table 1.

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Table 1

ſ				5	Synthetic E	Example N	10.		
	Compound	5	6	7	8	9	10	11	12
20	Solution A]				
25	#3 Sodium silicate (mole SiO ₂) Sodium hydroxide (mole NaOH)	273 g (1.0) 60 g (1.5)							
	Solution B								
30	Aluminum chloride • 6H ₂ O (mole Al ₂ O ₃) Cobalt chloride	145 g (0.3) 65 g	145 g (0.3)	145 g (0.3)	145 g (0.3)	145 g (0.3)	145 g (0.3) -	145 g (0.3)	145 g (0.3)
	(mole CoO) Nickel chloride*6H₂O (mole NiO)	(0.5)	120 g (0.5)	-	-	-	-	-	-
35	Ferric nitrate *9H ₂ O (mole FeO)	-	-	200 g (0.5)	-	-	-	-	-
	Copper chloride (mole CuO)	-	-	-	67 g (0,5)		-	•	•
	Titanium sulfate (mole TiO₂)	-	-	-	-	120 g (0.5)	-	•	
40	Barium chloride • 2H ₂ O (mole BaO)	-	-	-	-	-	122 g (0.5)	-	•
	Tin sulfate*2H₂O (mole SnO)	-	-	-	•	-	-	126 g (0.5)	-
45	Zirconium chloride (mole ZrO ₂)	-	-	-		-	-	-	116.5 g (0.5)

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Table 2

	•	sition Ratio o Components		
No.	SiO₂	MO _n	Al ₂ O ₃	BET Specific Surface Area
	(mole%)	(mole%)	(mole%)	(m²/g)
Synthetic Example 1	31	54	15	210
" 2	41	38	21	205
" 3	67	13	20	216
1 " 4	33	56	11	218
" 5	55	28	17	185
" 6	55	28	17	162
"7	55	28	17	158
" 8	55	28	17	163
" 9	55	28	17	171
" 10	55	28	17	160
" 11	55	28	17	154
" 12	55	28	17	172

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Each of the aluminosilicates obtained in Synthesis Examples 1 to 12, activated charcoal manufactured by Mitsubishi Kasei K.K. (Diasorb G*6-10) (specific surface area 900 m²/g), and various oxidizing agents shown in Table 3 were mixed to prepare deodorizer compositions (Present Products 1 - 12).

These products were evaluated by the evaluation methods shown below, and the evaluation results thereof are shown in Table 3 together with those of Comparative Examples.

Deodorizing Effect Evaluation Method A

Meat, vegetable, and fish were placed in a 6-liter desiccator and left to stand for 2 weeks to, prepare an artificial green garbage having an unpleasant odor.

A 5 g amount of the deodorizer composition was wrapped in an nonwoven cloth and placed in a wide mouth bottle to provide a sample. Further, from the 6-liter desiccator, 10 ml of the head space gas of the artificial green garbage odor was introduced into the wide mouth bottle, and the odor strength over a lapse of time was evaluated organoleptically, according to the following standards.

Evaluation point	Evaluation standard
0	no odor
1	very slight odor
2	slight odor
3	pervasive odor
4	strong odor
5	very strong odor

The evaluation was conducted by an expert panel of 5 members. The evaluations made by the top and bottom members were ignored, and an average of the evaluations made by the remaining three members was obtained by rounding off.

	1		1										-
5		ts over	After 6 hrs	o .	0	0	0	0	0	0	0	0	0
5		uation result lapse of time	After 3 hrs		1	п	7	0	0		0	0	н
10		Evaluation results lapse of time	After 1 hr	м	ന	7	7	1	ਜ	2	 4	H	2
15			Formulation amount (I) (a)/(b)	97/3	20/20	8/92	90/10	5/56	3/97	80/20	55/45	07/09	92/8
20			Forr amol										
25	Table 3	ltion	ring agent)		t e	٠					id	te	Sodium p-toluenesulfon chloroamide
30	Tabl	Deodorizer Composition	Component (b) (oxidizing agent)	rsulfate	Potassium monopersulfate	arbonate	eroxide	persulfate	Magnesium.perphthalate	Potassium permanganate	Diperoxy dodecane diacid	Potassium monopersulfate	oluenesulfon
35		Deodor	Component	Ammonium persulfate	Potassium n	Sodium percarbonate	Hydrogen peroxide	Potassium persulfate	Magnesium	Potassium	Diperoxy d	Potassium	Sodium p-t
40			(a)			H		ĸn	4	Ŋ	vo	7	ω
45			Component (a)	Activated charcoal	•	Synthetic Example	•	•	•	•		•	•
				г .	7	ю	4	52	9	7	80	Q	10
50				Example (Present product)									
		- 1		1									

5		rs over	After 6 hrs	0	o .	0	0	7	м	ν. ·
		Evaluation results over lapse of time	After 3 hrs	0	0	H	0	m	m	Ю
10		Evaluati laps	After 1 hr	п	-	2	ч	4	7	'n
15			Formulation amount (1) (a)/(b)	10/90	30/70	1/66	99.5/0.5	100/0	0/100	1
20			Forr amot				66	ਜ		
25	Table 3 (Continued)	sition	Component (b) (oxidizing agent)				nurate			
30	Table 3	Deodorizer Composition	ıt (b) (oxid	rsulfate	Armonium persulfate	lioxide	Sodium dichloroisocyanurate		Calcium hypochlorite	' .
35		Deode	Componer	Sodium persulfate	Ammonium 1	Chlorine dioxide	Sodium die		Calcium hy	
40			t (a)	ر و	10	11	. 12	.		
45			Component (a)	Synthetic Example	•	.	•	Activated charcoal	ı	1
50				#	12	13	14	-	7	ĸ
				Example (Present	product)			Compar- ative	Example	
55		1								

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A mixture of 90 g of the aluminosilicate obtained in Synthesis Example 1 and 10 g of potassium persulfate was kneaded with 50 g of water, by a mortar, followed by drying at 80°C to obtain a cake. The cake was crushed and sifted through a sieve to obtain a deodorizer composition as a white powder of 4 to 8 mesh (Present Product 15).

The results of the evaluation of this composition are shown below in Table 4, together with the results of Example 3.

Example 3

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A mixture of 95 g of silica powder, 5 g of sodium percarbonate was kneaded with 50 g of water, by a mortar, followed by drying at 80°C to obtain a cake. The cake was crushed and sifted through a sieve to obtain a white powder of 4 to 8 mesh. The white powder and the aluminosilicate obtained in Synthesis Example 2 were mixed to a formulation ratio of 9:1 to obtain a deodorizer composition (Present Product 16).

The deodorizing effect was evaluated according to the evaluation method described in Example 1, and the results of the evaluations of Examples 2 and 3 are shown in Table 4.

Table 4

25		Deodorizer	composition		ation resul	ì
		Component (a)	Component (b) (oxidizing agent)	After 1 hr	After 3 hrs	After 6 hrs
30	Present product 15 " 16 Comparative Example 4 " 5	Synthetic Example 1 " 2 Activated charcoal -	Potassium persulfate Sodium percarbonate - -	1 1 4 5	0 0 3 5	0 0 2 5

Example 4

A 2 g amount of dried leaves of rosemary was crushed by a cutter, charged into a Soxhlet extractor, and subjected to hot extraction with an addition of 180 g of water for 15 hours, whereby a 10% rosemary extract was obtained.

A 100 g amount of the aluminosilicate obtained in Synthesis Example 1 was impregnated with the 10% Rosemary extract and dried at 80°C, and the cake thus obtained was sifted through a sieve of 8 to 16 mesh to obtain a deodorizer composition.

Example 5

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A 20 g amount of the whole grass of thyme was crushed, charged into a Soxhlet extractor, and subjected to hot water extraction with an addition of 180 g of water for 13 hours to obtain an extract. A deodorizer composition was obtained in the same manner as in Example 4.

Example 6

A 10 g amount of leaves of dried sage was crushed, charged into a Soxhlet extractor, and subjected to

hot extraction with an addition of 190 g of 50% ethanol for 15 hours to obtain a 5% sage extract. An amount of 100 g of the aluminosilicate obtained in Synthesis Example 2 was impregnated with 10 g of the 50% sage extract obtained by concentration by an evaporator, followed by drying at 110 °C. The cake obtained was sifted through a sieve of 8 to 16 mesh to obtain a deodorizer composition.

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The 10% rosemary extract obtained in Example 4 was sprayed onto 100 g of the aluminosilicate obtained in Synthesis Example 3 to obtain a powdery deodorizer composition.

Example 8

Example 7

The 10% rosemary extract obtained in Example 4 was sprayed onto activated charcoal (manufactured 15 by Mitsubishi Kasei K.K.; Diasorb G-6-10) to obtain a powdery deodorizer composition.

Example 9

The sage extract obtained in Example 6 was sprayed onto spherical silica to obtain a deodorizer composition.

Example 10

A 30 g amount of pine leaves was crushed, charged into a Soxhlet extractor, and subjected to hot extraction with an addition of 150 g of acetone for 10 hours to obtain an extract. A deodorizer composition was the obtained by the same treatment as used in Example 4.

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Evaluation of the Deodorizing Effect (Examples 4 - 10)

The deodorizer compositions of Examples 4 to 10 were evaluated according to the above-mentioned evaluation method A. 35

The results are shown in Table 5.

Table 5

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		Deoc	lorizing	effect
	Sample	1 hr	3 hrs	6 hrs
Present product	Example 4 " 5 " 6 " 7 " 8 " 9 " 10	1 2 1 1 3 3	0 1 0 0 1 1	0 0 0 0 0
Comparative Example	6 (10% Rosemary extract) 7 (Activated charcoal) 8 (50% Sage extract) 9 (Blank)	4 4 4 5	4 3 4 5	3 3 3 5

Examples 11 - 18

The aluminosilicates obtained in Synthesis Examples 1 to 3, activated charcoal manufactured by Mitsubishi Kasei K.K. (Diasorb G-6-10), and various sterilizers shown in Table 6 were mixed to prepare deodorizer compositions.

These were evaluated according to the following evaluation method (method B), and the evaluation results thereof are shown in Table 6 together with those of Comparative Examples.

Deodorizing Effect Evaluation Method B

Meat, vegetables, fish, were placed in a 6-liter desiccator and left to stand for one week, to prepare an artificial green garbage. An amount of 30 g of the green garbage prepared was placed into a wide mouth bottle of 1.8-liter, 10 g of the deodorizer composition sprayed thereon, and the odor intensity over a lapse of time was evaluated organoleptically according to the following standards.

Evaluation point	Evaluation standard
0	no odor very slight odor
2	slight odor
3 4	pervasive odor strong odor
5	very strong odor

Evaluation was conducted by an expert panel of 5 members. The top and bottom members were ignored, and an average of the evaluations of the remaining three members was obtained by rounding off.

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			After 2 days	ю	m	2	4	٠ ١
5		Evaluation results over lapse of time	After After After 6 hrs 1 day 2 day	7	e	2	m.	'n
10		uation result: lapse of time	After 6 hrs	7	ო	8	ю	ĸ
		valuati laps	After 3 hrs	m	m	m	m	2
15		ы	After 1 hr	4	4	7	4	S
20			Formulation ratio (a)/(b)	100/0	0/100	99.995/0.005	100/0	1
25	inued)						٠	
30	Table 6 (Continued)	Deodorizer Composition	Component (b) (Germicide)	ı	oride	oate	ı	ı
35		Deodorize	Component (Benzalkonium chloride	Methyl p-oxybenzoate		
			(a)		—	Σ	der	
45			Component (a)	Activated charcoal	ı	Activated charcoal	Silica powder	ı
50				10	11	12	13	14
55				Compar- ative	Example			

			After 2 days	o	0	0	0	0	0	0	0
5		lts over	After 1 day	0	0	0	0	0	0	0	0
10		Evaluation results lapse of time	After 6 hrs	0	0	0	0	0	0	0	0
		raluatio lapse	After 3 hrs	Ħ	ч	-	0	н	0	~ 1	н
15		ы	After 1 hr	m	2	2	ч	7	7	м	2
20	,		Formulation ratio (a)/(b)	99/1	1.0/6.66	50/50	90/10	97/3	95/5	70/30	97/3
25											
30	Table 6	Deodorizer Composition	Component (b) (Germicide)	ride		nzimidazole	.)-1- 191)urea	.2'-hydroxy	10	P.	1 dehyde
35		Deodorizer	Component (b	Benzalkonium chloride	o-Phenylphenol	2-(4-Thiazolyl)benzimidazole	3-(4-Chlorophenyl)-1- (3,4-dichlorophenyl)urea	4,4'-Trichloro-2'-hydroxy phenyl ether	p-Chloro-m-xylenol	Dehydroacetic acid	α-Bromocinnamic aldehyde
40				Benza	o-Phe	2-(4-	3-(4-	2,4,4 diphe	p-Ch1	Dehyd	a-Bro
			(a)				~	7	ബ	7	7
45			Component (a)	Activated charcoal	•	•	Synthetic Example	•	-	=	•
50				11	12	13	14	15	16	17	18
55				Example (Present							
		ı		1							

A 100 g amount of the aluminosilicate obtained in Synthesis Example 1 was impregnated with 50 g of a 0.2% ethyl alcohol solution of 2-(4-thiazolyl)benzimidazole, and dried at 80°C for one hour to obtain a powdery deodorizer composition.

Example 20

Onto 50 g of activated charcoal (manufactured by Mitsubishi Kasel, Diasorb G-6-10) was sprayed 10 g of an aqueous 30% benzalkonium chloride, to obtain a powdery deodorizer composition.

The deodorizing effect of the deodorizer compositions of Examples 19 and 20, was evaluated according to the method B. The results are shown in Table 7.

	Doodori	Deederizer composition	Eva	luation re	sults over	Evaluation results over lapse of time	time
	ionopo	zoi composition					
	Component (a)	Commonent (b) (Germicide) After 1 After 3 After 6 After 1 After 2	After 1	After 3	After 6	After 1	After 2
	(a) monodino		È	hrs	hrs	day	days
							,
0 0	Synthetic Example 1	Swithestic Example 1 2-(4-Thiazolyl)benzimidazole	_	0	0	0	0
Example 13	cylinder Example:		,	,	_	_	_
	Activated charcoal	Benzalkonium chloride	77	-	>	>	>
03			_	ď	~	2	က
Comparative Example 15 Activated charcoal	Activated charcoal	ı	-		J 1		Ų
		•	2	ß	c.	ດ	0
	•		ļ				

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A 100 g amount of the aluminosilicate obtained in Synthesis Example 3 was impregnated with 30 g of 10% ethyl alcohol solution of o-phenylphenol, and dried at 80°C for 30 minutes to obtain a powdery deodorizer composition.

When the deodorizer composition was sprayed into a polymer bucket and a triangular sink-corner container for garbage, containing green garbage, the odor generated from the green garbage or remaining foods was eliminated.

Examples 22 - 28

Samples of each of the wastewaters shown below were placed, to an amount of one liter, in a 1.8-liter glass bottle.

Wastewater-A: washed wastewater from pig sty (feeding 87 pigs)

Wastewater-B: wastewater from cleaning tank of a standard family having 4 members.

Then 10 g of the deodorizer composition shown in Table 5 was added thereto, and the odor intensity over a lapse of time was organoleptically evaluated. The evaluation standards were made according to method B.

The deodorizer composition was a powdery product obtained by impregnating the aluminosilicate obtained in each Synthesis Example with an ethanolic solution of the sterilizer, followed by drying at 80°C for one hour, and supplemented with 1% of sterilizer.

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		I able 8						
	Deodori	Deodorizer composition	Treated waser	Eva	luation re	sults over	Evaluation results over lapse of time	time
	Component (a)	Component (b) (Germicide)		After 1	After 3	After 6	1	After 2
	•			훋	hrs	hrs	day	days
Example 22	Synthetic Example 2	o-chloro-m-xylenol	A	8	-	0	0	0
"23	. 3	o-phenylphenol	0	8	-	0	0	0
" 24		2-(4-thiazolyl)benzimidazole	œ	2	-	0	0	0
25	- 2	Dehydroacetic acid	ω	က	-	0	0	0
26	- 2	o-phenylphenol	⋖	2	-	0	0	0
" 27	<u>.</u>	Benzalkonium chloride	۷	က	-	0	0	0
28	2	Butyl m-cresol	⋖	7	-	0	0	0
Comparative Example 17	Silica		ω	4	က	က	ო	4
18		Dehydroacetic acid	œ	4	က	က	ო	င
0 0	Activated charcoal	,	⋖	4	က	က	က	4
2								

Examples 29 - 31

Onto 10 g of the aluminosilicate obtained in Synthesis Examples 1 - 3, 2 g of the 10% rosemary extract was sprayed and dried at 80°C for 3 hours, and the sample obtained was mixed with 0.3 g of a sterilizer to obtain a deodorizer composition.

The evaluations were made according to the evaluation method (method B), and the evaluation results are shown in Table 9 together with Comparative Examples.

		_	D DICIG					
	800	Deodorizer composition	osition	Eval	uation re	sults over	Evaluation results over lapse of time	time
	חפר	ממווקה הפווחת						
		plant extract	Germicide (b) (iii)	After 1	After 3	After 6	After 1	After 1 After 3 After 6 After 1 After 2
	component (a)	(5)(6)		È	hrs	hrs	day	days
		//					,	,
-	Posemany Bosemany	Boseman	o-nhenvlphenol	7	-	0	0	>
Example 29	Symmenc Evample of	, minocon		_		-	-	_
. 30		=	Benzalkonium chloride	V	_	>	>) (
6				•	-	0	0	0
.31	2"	=	Denydroacetic acid	J	. ,	. (c
		•	Benzalkonium chloride	4	က	.7	N	ာ
Comparative example 20				•	ď	~	67	ന
. 21	Activated charcoal		•	+	,	,	,	
			pioc citodocal.	~	•	۰	~	2
6		Rosemary		>	J	_		

Claims

- 1. A deodorizer composition comprising:
 - (a) an activated charcoal, a silica gel, a clay mineral, or an aluminosilicate having the following composition, in terms of oxides, represented by the composition ratio of the three components:

SiO₂: 5 to 80 mole%; Monz: 5 to 65 mole%;

10 Al₂O₃: 1 to 60 mole%

- wherein M represents at least one metal selected from the group consisting of zinc, copper, silver, cobalt, nickel, iron, titanium, barium, tin and zirconium, and n represents a valence of metal; and
- (b) at least one component selected from the group consisting of (i) oxidizing agents, (ii) plant extracts, and (iii) germicides.
- 2. A deodorizer composition as claimed in claim 1, wherein the weight ratio of the components (a)/(b) is from 1/99 to 99.99/0.01.
- 3. A deodorizer composition as claimed in claim 1 or 2, wherein said oxidizing agent is an organic peracid or a salt thereof, a percarbonate, or a persulfate.
- 4. A deodorizer composition according to claim 1 or 2, wherein the plant extract is selected from among red algae, brown algae, gymnosperms and angiosperms.
- 5. A deodorizer composition as claimed in claim 1 or 2, wherein the germicide is a sublimable substance.
- 6. A deodorizer composition as claimed in claim 1 or 2, wherein the germicide is p-chloro-m-xylenol, butyl-m-cresol, o-phenylphenol or α -bromocinnamic aldehyde.

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EUROPEAN SEARCH REPORT

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	Citation of document with indi	cation, where appropriate.	Relevant	CLASSIFICATION OF THE
ategory	of relevant pass		to claim	APPLICATION (Int. Cl.5)
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X	FR-A-1 428 096 (AME * Examples 1-3 *	RICAN CYANAMID CO.)	1,3	
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A	WO-A-8 101 643 (MOL SYSTEMS)	ECULAR SIEVE		TECHNICAL FIELDS SEARCHED (Int. Cl.5) A 61 L
	The present search report has be	en drawn up for all claims		
Place of search THE HAGUE CATEGORY OF CITED DOCUMENTS Date of completion of the search D6-04-1990 T: theory or p E: earlier pate			PINOSA Y CARRETERO	

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